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Polyisocyanurate foam.

- The present invention provides for a polyisocyanurate foam comprising the reaction product in the presence of a blowing agent and catalyst of
 - (a) an isocyanate group-terminated quasiprepolymer prepared by reacting an organic polyisocyanate with-an organic compound having at least two active hydrogen atoms, and
 - (b) a minor amount of polyol component, characterized in that the polyol component comprises
 - (i) a polyester polyol having a free glycol content of less that about 7 percent by weight of the polyester polyol or
 - (ii) a mixture of the low free glycol-containing polyester polyol with at least one other polyol, the total free glycol content of mixture (b) being less that about 7 percent by weight of the polyester polyol.

Also disclosed is a polyur than foam and laminate products utilizing the polyisocyanate and polyurethane foams.



EUROPEAN SEARCH REPORT

EP 90 10 6790

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x	EP-A-0 278 412 (BASF) * page 6, lines 18-29; page 13 claims *; US - A - 4791148 (ca	s, table V, examples 21-24;	1-7,9	C 08 G 18/42 C 08 G 18/66 //
Y	Claims , 000 A 1101110 (St		1-10	(C 08 G 18/10 C 08 G 101:00)
X	US-A-4 636 530 (T. NARAY) * column 5, lines 3-65; columi	AN) n 9, table II; claims 1-3,8-12 *	1-4,6	3,7
Y	_		1-10	
Y	US-A-4 237 238 (D.T. DEGU column 2, lines 4-67; column 25-31,42-48; claims 1-6	JISEPPI et al.) n 3, lines 53-60; column 4, lines	1-10	
Y	US-A-4 758 607 (R.K. HALL * abstract; column 9, lines 27 column 15, line 9 *	MARK et al.) -55; column 14, line 59 -	1-10	
Y	US-A-4 496 625 (S.C. SNID * abstract; claims 1,7-10 *	ER et al.)	1-10	
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	A: technological background O: non-written disclosure P: Intermediate document T: theory or principle underlying the it	do	mber of	f the same patent family, corresponding

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- The present invention provides for a polyisocyanurate foam comprising the reaction product in the presence of a blowing agent and catalyst of
 - (a) an isocyanate group-terminated quasi-prepolymer prepared by reacting an organic polyisocyanate with an organic compound having at least two active hydrogen atoms, and
 - (b) a minor amount of polyol component, characterized in that the polyol component comprises
 - (i) a polyester polyol having a free glycol content of less that about 7 percent by weight of the polyester polyol or
- (ii) a mixture of the low free glycol-containing polyester polyol with at least one other polyol, the total free glycol content of mixture (b) being less that about 7 percent by weight of the polyester polyol. Also disclosed is a polyurethane foam and laminate products utilizing the polyisocyanate and polyurethane foams.

This invention r lates to the polyisocyanurate and polyurethane foams characterized by improved insulating prop rties. Mor particularly, the invention relates to an improvement in the foams insulating capacity by reacting a polyisocyanate prepolymer and polyester polyol containing low free glycol.

The preparation of foams characterized by isocyanurate and urethane linkages is well known in the art. Generally, these foams are prepared by reacting an organic polyisocyanate with a polyol in the presence of a blowing agent and a catalyst(s). Polyester polyols of many types can be used as the polyol components in the production of these foams.

Because of the relatively high cost of various typical polyols useful in making cellular polymers and because an inexpensive source of polyols is now available from such sources as crude dimethyl terephthalate oxidate residues, recycled poly (ethylene terephthalates), and various other scrap polyester sources, efforts have heretofore been directed to processes for converting these materials to commercially useful polyester polyols. These efforts have led to a number of polyester polyols found to be highly suitable in producing foams of good quality. However, there is still a search for improvements in these polyols and the ways of using them in foam manufacture.

It is also known that polyols, including certain aromatic polyester polyols, can be reacted with isocyanates to form isocyanate-terminated quasi-prepolymers which in turn can be used in the formation of polyurethane and/or polyisocyanurate foams, as disclosed, e.g., in U. S. Patent No. 4,791,148.

A most important characteristic of rigid foams is their insulating value which is expressed by the k- and aged k- factor. The k-factor is a measurement of the insulating properties (thermoconductivity) and the aged k-factor indicates insulating properties over time. Foams prepared from the above-noted quasi-prepolymers of U. S. Patent No. 4,791,148 exhibit low initial thermal conductivity values and maintain low thermal conductivity values after prolonged periods of use. It would be most beneficial to provide still further improved formulations for the production of rigid foams having a combination of advantageous properties, including especially good insulating properties, as reflected in low initial thermal conductivity values and low thermal conductivity values after prolonged periods of use.

The present invention relates to the production of improved rigid plastic foams, especially polyisocyanurate and polyurethane foams, having a combination of advantageous properties, including excellent thermal insulative properties, reduced friability, and good dimensional stability, thermal resistance, and compressive strength, and an improved method of producing the foams.

The present invention also is directed to the production of closed cell foam materials which can be used in building panels which are highly insulating, thermally resistant, low in friability, and soundproof.

The present invention also concerns the production of a polyester polyol-containing composition for use in the preparation of rigid plastic foams having a combination of advantageous properties, especially an improved insulating efficiency.

The present invention relates to the combined use of a polyisocyanate prepolymer and a polyester polyol composition containing a low free glycol content for the formation of rigid plastic foams having improved insulating properties. While the polyisocyanate prepolymer and low free glycol-containing polyester polyol compositions individually contribute to the formation of low k-factor foams, it has now been surprisingly discovered that their combined use results in the production of foams having lower k-factors than foams made with either composition alone.

More particularly, the invention relates to a polyisocyanurate foam comprising the reaction product in the presence of a blowing agent and catalyst of

- (a) an isocyanate group-terminated quasi-prepolymer prepared by reacting an organic polyisocyanate with an organic compound having at least two active hydrogen atoms, and
 - (b) a minor amount of a polyol component, characterized in that the polyol component comprises
- (i) a polyester polyol having a free glycol content of less than about 7 percent by weight of the polyester polyol or
- a mixture of the low free glycol-containing polyester polyol with at least one other polyol, the total free glycel content of mixture (b) being less than about 7 percent by weight of the polyester polyol.

The invention further particularly relates to a polyurethane foam comprising the reaction product in the presence of a blowing agent and a catalyst of

- (a) an isocyanate group-terminated quasi-prepolymer prepared by reacting an organic polyisocyanate with an organic compound having at least two active hydrogen atoms, and
- (b) a polyol component, characterized in that the polyol component comprises a polyol mixture containing a polyester polyol having a free glycol content of less than about 7 percent by weight of the polyester polyol, the amount of the low free glycol-containing polyest r polyol in the polyol mixture being sufficient to reduce the aged k-factor of the resultant foam to below the aged k-factor of the corresponding foam prepared from the same foam-forming composition and having the same indix accept that the polyol

component does not include the low free glycol-containing polyester polyol.

Further provided are laminate products utilizing the polyisocyanurate and polyurethane foams.

The polyisocyanate prepolymers (quasi- prepolymers) may be prepared by reacting an excess of an organic polyisocyanate or a mixture of polyisocyanates with a minor amount of at least one active hydrogen containing compound. The low free glycol-containing polyester polyol compositions of the invention may be prepared by reacting a polycarboxylic acid and/or acid derivative with a polyol component comprising at least one glycol compound. The improved foams of the invention advantageously comprise polyisocyanurate and polyurethane foams prepared by reacting together the isocyanate-terminated quasiprepolymer and the low free glycol-containing polyester polyol composition, and optionally other polyhydroxyl compounds in the presence of a blowing agent(s). The reaction may be carried out in the presence of catalysts, auxiliaries and additives as required (e.g., a surfactant).

The isocyanate-terminated quasi-prepolymers of the invention are prepared by reacting together one or more organic isocyanates with an organic compound having at least two active hydrogen atoms as determined by the well-known Zerewitinoff test, as described by Kohler in Journal of the American Chemical Society, 40, 318 (1927). The reaction takes advantage of the reactivity of the isocyanate groups and builds up multiple molecular units of higher molecular weight. Chain building compounds containing active hydrogen atoms which are particularly useful are polyester and polyether polyols, especially polyester polyols and mixtures of polyester and polyether polyols. The chain builders are used in predetermined amount sufficient to produce reactive polyisocyanate prepolymers of the desired equivalent weight (e.g., 140 to 250) and significant excess isocyanate functionality. Generally, the quasi-prepolymers have a free isocyanate content of from 16.8 % to 30.0 %, preferably 24.0 % to 29.0 %, by weight.

The preparation of the quasi-prepolymer may be exemplified with reference to that derived from a polyester polyol. The preparation is accomplished simply by reacting the isocyanate and polyester polyol in a suitable reaction vessel, generally in an inert (nitrogen) atmosphere. The Isocyanate is preferably heated 25 to approximately 50° to 100° C. before addition of polyester polyol, which is added gradually with stirring over a period of time generally between 15 minutes and several hours. A modest exotherm is allowed to develop and the temperature is increased to approximately 90° C. and held there until the desired NCO content is reached, after which the product is discharged.

The quantity of polyester polyol which is reacted with the isocyanate may vary, but generally is not less than 2 percent and not more than 30 percent relative to the weight of the final quasi-prepolymer, and preferably 20 percent or less. The stoichlometry must be such that sufficient equivalents of isocyanate are present to insure that the finished quasi-prepolymer will be isocyanate-terminated, as is well known to those skilled in the art. Examples of analogous formulations and their stoichiometrics may be found in POLYURETHANES: CHEMISTRY AND TECHNOLOGY, volumes I and II, by Saunders and Frisch, published by Wiley-Interscience. Catalysis is generally not required to form the quasi-prepolymers, and it is undesirable to include non-removable catalysts whose residual catalytic effect would cause further polymerization leading to undesirable increases in viscosity during storage. However, catalysts which do not promote the reaction of isocyanate groups with each other may be used, especially when the quasiprepolymer is to be foamed within a short time.

Catalysts suitable for the preparation of the isocyanate-terminated prepolymers are, for example, the various alkyl tin carboxylates and tin carboxylates. Preferably used when catalysis is desired, for example, are tin-(II) octoate, and dibutyl tin dilaurate. When catalysts are used, they may be present in amounts at from 0.005 to 2 percent by weight, preferably from 0.01 to 0.1 percent by weight. As indicated previously, preferably no catalyst is utilized.

The organic isocyanates to be reacted with the chain builders have a functionality of two or higher. For . example, organic diisocyanates, polyisocyanates, or mixtures thereof may be used successfully. The organic isocyanates may be aliphatic, cycloaliphatic, alicyclic, aromatic or aromatic aliphatic isocyanates. The various isocyanates suitable for the preparation of the quasi-prepolymers of the invention are well known to those skilled in the art.

Among the many isocyanates suitable for the practice of the subject invention are, for example, aliphatic isocyanates such as tetramethylene, hexamethylene, octamethylene and decamethylene diisocyanates, and their alkyl substituted homologs; cycloaliphatic isocyanates such as 1,2-, 1,3- and 1,4cyclohexane diisocyanates, 2,4- and 2,6-methylcyclohexane diisocyanates, 4,4'- and 2,4'-dicyclohexyldiisocyanates, 4,4'- and 2,4'-dicyclohexylmethane diisocyanates, 1,3,5-cyclohexane triisocyanates, and saturated (hydrogenated) polymethylenepolyphenylen polyisocyanates; alicyclic isocyanates such as Isocyanatom thylcyclohexane isocyanates, isocyanatoethylcyclohexane isocyanates, bis(isocyanatomethyl)cyclohexane diisocyanates, 4,4 - and 2,4 -bis(isocyanatomethyl) dicyclohexane, and isophorone diisocyanate; aromatic isocyanates such as 1,2-, 1,3-, and 1,4-phenylene diisocyanates, 2,4- and 2,6-toluene diisocyanate, 2,4'-, 4,4'-and 2,2-biphenyl diisocyanates, 2,2'-, 2,4'- and 4,4'-diphenylmethane diisocyanates, and polymethylenepolyphenylenepolyisocyanates (polymeric MDI); and aromatic aliphatic isocyanates such as 1,2-, 1,3-, and 1,4-xylylene dilsocyanates.

Organic isocyanates containing heteroatons may also be utilized, for example those derived from melamine. Modified polyisocyanates, such as carbodiimide or isocyanurate can also be employed. Liquid carbodiimide group- and/or isocyanurate ring-containing polyisocyanates having isocyanate contents from 15 to 33.6 percent by weight, preferably from 21 to 31 percent by weight, are also effective, for example, those based on 4.4 -, 2.4 -, and/or 2.2 -diphenylmethane diisocyanate and/or 2.4- and/or 2.6-toluene diisocyanate, and preferably 2,4- and 2,6-toluene diisocyanate and the corresponding isomer mixtures, 4,4-, 2,4, and 2.2 diphenylmethane diisocyanates as well as the corresponding isomer mixtures, for example, mixtures of 4,4'- and 2,4'-diphenylmethane diisocyanates, mixtures of diphenylmethane diisocyanates and polyphenyl polymethylene polyisocyanates (polymeric MDI), and mixtures of toluene diisocyanates and polymeric MDI. Preferred, however, are the aromatic diisocyanates and polyisocyanates. Particularly preferred are 2,4-, and 2,6-toluene diisocyanate and mixtures thereof (TDI), 2,4'-, 2,2'- and 4,4'-diphenylmethane diisocyanate (MDI), polymethylenepolyphenylenepolyisocyanates (polymeric MDI), and mixtures of the above preferred isocyanates.

Most particularly preferred are the polymeric MDIs.

Suitable active hydrogen-containing groups as determined by the Zerewitinoff method which are reactive with an isocyanate group include -OH, -NH-, -COOH, and -SH. Examples of suitable types of organic compounds containing at least two active hydrogen-containing groups which are reactive with an isocyanate group are hydroxyl-terminated polyesters, polyalkylene polyether polyols, hydroxy-terminated polyurethane polymers, polyhydric polythioethers, alkylene oxide adducts of phosphorus-containing acids, polyacetals, aliphatic polyols, aliphatic thiols including alkane, alkene and alkyne thiols having two or more -SH groups; diamines including both aromatic, aliphatic and heterocyclic diamines, as well as mixtures thereof. Compounds which contain two or more different groups within the above-defined classes may also be used in accordance with the process of the present invention such as, for example, amino alcohols which contain an amino group and a hydroxyl group. Also, compounds may be used which contain one -SH group as well as those which contain an amino group and a -SH group.

In a preferred embodiment of the invention, polyester polyols are used as chain builders in producing the isocyanate terminated quasi-prepolymers. However, whether prepared from a polyester polyol or some other chain builder, the quasi-prepolymers are polymerized in the presence of a specific polyol, viz., a polyester polyol having a low free glycol content, to form the inventive polyisocyanurate and/or polyurethane foams which exhibit low initial thermal conductivity values and maintain low thermal conductivity values after prolonged periods of use.

The polyester polyols themselves can be prepared by known procedures from a polycarboxylic acid component comprising a polycarboxylic acid or acid derivative, such as an anhydride or ester of the polycarboxylic acid, and any polyol component, the polyol component comprising a glycol(s) or a glycolcontaining mixture of polyois in the production of the low free glycol-containing polyester polyoi. The polyacid and/or polyol components may, of course, be used as mixtures of two or more compounds in the preparation of the polyester polyols. Particularly suitable polyester polyols for use in preparing the quasiprepolymers and for reaction with the quasi-prepolymers in foam production are aromatic polyester polyols containing phthalic acid residues.

The production of the polyester polyols is accomplished by simply reacting the polycarboxylic acid or acid derivative with the polyol component in a known manner until the hydroxyl and acid values of the reaction mixture fall in the desired range. The reaction typically is performed at temperatures from about 150°C to 250°O for a period from about 1 to 10 hours. The reaction can be carried out as a batch process or continuously. When present, excess glycol can be distilled from the reaction mixture during and/or after the reaction, such as in the preparation of the low free glycol-containing polyester polyols. Normally, an effective amount of catalyst is added to promote the present reaction. Any conventional esterification or transesterification catalyst (single compound or mixture of compounds) can be used. Suitable catalysts include organotin compounds, particularly tin compounds of carboxylic acids, such as stannous octoate, stannous oleate, stannous acetate, stannous laurate, dibutyl tin dilaurate, and other such tin salts. Additional suitable metal catalysts include tetraisopropyl titanate, and other such titanate salts, and the like.

The polycarboxylic acid component may be aliphatic, cycloallphatic, aromatic and/or heterocyclic and 55 may optionally be substituted, for example, by halogen atoms, and/or may be unsaturated. Examples of suitabl carboxylic acids and derivatives thereof for the pr paration of th polyester polyols include: oxalic acid; malonic acid; succinic acid; glutaric acid; adipic acid; pimelic acid; suberic acid; az laic acid; sebacic acid; phthalic acid; isophthalic acid; trimellitic acid; ter phthalic acid; phthalic acid anhydride; tetrahydrophthalic acid anhydride; pyromellitic dianhydride; hexahydrophthalic acid anhydride; t trachlorophthalic acid anhydride; endomethylene tetrahydrophthalic acid anhydride; glutaric acid anhydrid; maleic acid; mal ic acid anhydride; fumaric acid; dibasic and tribasic unsaturated fatty acids optionally mixed with monobasic unsaturated fatty acids, such as oleic acid; terephthalic acid dimethyl ester and terephthalic acid-bisglycol ester.

ester.

Polyester polyols whose acid component advantageously comprises at least about 30% by weight of phthalic acid residues are particularly useful. By phthalic acid residue is meant the group

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While the aromatic polyester polyols can be prepared from substantially pure reactant materials, more complex ingredients are advantageously used, such as the side-stream, waste or scrap residues from the manufacture of phthalic acid, terephthalic acid, dimethyl terephthalate, polyethylene terephthalate, and the like. Particularly suitable compositions containing phthalic acid residues for use in the invention are (a) ester-containing by-products from the manufacture of dimethyl terephthalate, (b) scrap polyalkylene terephthalates. (c) phthalic anhydride, (d) residues from the manufacture of phthalic acid or phthalic anhydride, (e) terephthalic acid, (f) residues from the manufacture of terephthalic acid, (g) isophthalic acid, (h) trimellitic anhydride, (i) phthalic acid, and (j) combinations thereof. These compositions may be converted by reaction with the polyols of the invention to polyester polyols through conventional transesterification or esterification

procedures.

A preferred polycarboxylic acid component for use in the preparation of the aromatic polyester polyols is phthalic anhydride. This component can be replaced by phthalic acid or a phthalic anhydride bottoms composition, a phthalic anhydride crude composition, or a phthalic anhydride light ends composition, as such compositions are defined in U. S. Patent No. 4,529,744.

Other preferred materials containing phthalic acid residues are polyalkylene terephthalates, especially polyethylene terephthalate (PET), residues or scraps.

Still other preferred residues are DMT process residues, which are waste or scrap residues from the manufacture of dimethyl terephthalate (DMT). The term "DMT process residue" refers to the purged residue which is obtained during the manufacture of DMT in which p-xylene is converted through oxidation and esterification with methanol to the desired product in a reaction mixture along with a complex mixture of by-products. The desired DMT and the volatile methyl p-toluate by-product are removed from the reaction mixture by distillation leaving a residue. The DMT and methyl p-toluate are separated, the DMT is recovered and methyl p-toluate is recycled for oxidation. The residue which remains can be directly purged from the process or a portion of the residue can be recycled for oxidation and the remainder diverted from the process, or, if desired, the residue can be processed further, as, for example, by distillation, heat treatment and/or methanolysis to recover useful constituents which might otherwise be lost, prior to purging the residue from the system. The residue which is finally purged from the process, either with or without additional processing, is herein called DMT process residue.

These DMT process residues may contain DMT, substituted benzenes, polycarbomethoxy diphenyls, benzyl esters of the toluate family, dicarbomethoxy fluorenone, carbomethoxy benzocoumarins and carbomethoxy polyphenols. Hercules, Inc., Wilmington, Del., sells DMT process residues under the trademark Terates 101. DMT process residues having a different composition but still containing the aromatic esters and acids are also sold by DuPont and others. The DMT process residues to be transesterified in accordance with the present invention preferably have a functionality at least slightly greater than 2.

Such suitable residues include those disclosed in U.S. Patent Nos. 3,647,759, 4,411,949, and 4,714,717. The polyester polyols which are used as chain builders in producing the isocyanate terminated quasi-prepolymers may be prepared from the above described polycarboxylic acid components and any polyol component. The polyols can b aliphatic, cycloaliphatic, aromatic and/or heterocyclic. Low mol cular weight aliphatic polyhydric alcohols, such as aliphatic dihydric alcohols having no more than about 20 carbon atoms are highly satisfactory. The polyols optionally may include substituents which are linert in the r action, for exampl, chlorine and bromine substituents, and/or may be unsaturated. Suitable amino

alcohols, such as, for exampl , monoethanolamine, diethanolamine, triethanolamine, or the like may also be used. Moreov r, the polycarboxylic acid(s) may be condensed with a mixture of polyhydric alcohols and amino alcohols.

Examples of suitable polyhydric alcohols include: ethylene glycol; propylene glycol-(1,2) and -(1,3); butylene glycol-(1,4) and -(2,3); hexane diol-(1,6); octane diol-(1,8); neopentyl glycol; 1,4-bishydroxymethyl cyclohexane; 2-methyl-1,3-propane diol; glycerin; trimethylolpropane; trimethylolethane; hexane triol-(1,2,6); butane triol-(1.2.4); pentaerythritol; quinitol; mannitol; sorbitol; formitol; α-methyl-glucoside; diethylene glycol; triethylene glycol; tetraethylene glycol and higher polyethylene glycols; dipropylene glycol and higher polypropylene glycols as well as dibutylene glycol and higher polybutylene glycols. Especially suitable polyols are alkylene glycols and oxyalkylene glycols, such as ethylene glycol, diethylene glycol, dipropylene glycol, triethylene glycol, tripropylene glycol, tetraethylene glycol, tetrapropylene glycol, trimethylene glycol and tetramethylene glycol, and 1,4-cyclohexanedimethanol(1,4-bis-hydroxymethylcyclohexane).

The term "polyester polyol" as used in this specification and claims includes any minor amounts of unreacted polyol remaining after the preparation of the polyester polyol and/or unesterified polyol added after the preparation.

These prepolymer-building polyester polyols advantageously have an average functionality of about 1.8 to 8, preferably about 1.8 to 5, and more preferably about 2 to 2.5. Their hydroxyl number values generally fall within a range of about 15 to 750, preferably about 30 to 550, and more preferably about 100 to 550.

The quasi-prepolymer may be produced from commercially available polyester polyols. Examples of such polyester polyols are those derived from PET scrap and available under the designation Chardol 170, 570, 571 and 572 from Chardonol and Freol 30-2150 from Freeman Chemical. Examples of suitable DMT derived polyester polyols are Terate® 202, 203, and 204 polyols, which are available from Hercules Incorporated. Phthalic anhydride derived-polyester polyols are commercially available under the designation Pluracol® polyol 9118 from BASF Corporation, and Stepanol PS-2002, PS-2502A, PS-2502A, PS-2502, PS-2522, PS-2852, PS-2552, and PS-3152 from Stepan Company. NCO-capped prepolymers prepared from the phthalic anhydride derived polyester polyois, and the foams prepared therefrom constitute preferred embodiments of the present invention.

In the preparation of the foams of the present invention, the quasi-prepolymer is reacted with a low free glycol-containing polyester polyol whose free glycol content is sufficiently low to bring about significant improvement in foam insulation value. Typically, the free glycol content is less than about 7, preferably less than about 6, and more preferably less than about 5, percent by weight of the total polyester polyol

The glycols which are reacted with the above-described polycarboxylic acid components in producing 35 the low free glycol-containing polyester polyols may contain heteroatoms (e.g., thiodiglycol) or may be composed solely of carbon, hydrogen, and oxygen. They are advantageously simple glycols of the general formula C_nH_{2n}(OH)₂ or polyglycols distinguished by intervening ether linkages in the hydrocarbon chain, as represented by the general formula $C_nH_{2n}O_x(OH)_2$. In a preferred embodiment of the invention, the glycol is a low molecular weight aliphatic diol of the generic formula:

HO-R-OH wherein R is a divalent radical selected from the group consisting of:

(a) alkylene radicals each containing from 2 through 6 carbon atoms, and

(b) radicals of the formula:

-(R1O)m-R1-

wherein R1 is an alkylene radical containing from 2 through 6 carbon atoms, and m is an integer of from 1 through 4, and - -

(c) mixtures thereof.

Besides the glycols, any suitable polyhydric alcohol may be used in preparing the free glycolcontaining polyester polyols. These additional polyols can be aliphatic, cycloaliphatic, aromatic and/or heterocyclic, and are preferably selected from the group consisting of diols, triols and tetrols. The polyols optionally may include substituents which are inert in the reaction, for example, chlorine and bromine substituents, and/or may be unsaturated. Suitable amino alcohols, such as, for example, monoethanolamine, diethanolamine, triethanolamine, or the like, may also be used. These additional polyols are used in amounts which do not detract from the enhanced insulating properties stemming from the low free glycol content of the polyester polyols.

A preferred low free glycol-containing poly ster polyol of the invention is prepared by reacting a polycarboxylic acid and/or acid derivative, especially one containing a phthalic acid r sidue (e.g., phthalic anhydride), with an alkylene glycol having the formula

wherein R² is hydrogen or lower alkyl of one to four carbon atoms, preferably hydrogen or methyl, and z is from 1 to 5, preferably 1 to 3, and more preferably 1 to 2. Examples of the alkylene glycols are ethylene glycol, propylene glycol, diethylene glycol, dipropylene glycol, triethylene glycol, tetraethylene glycol, tripropylene glycols and tetrapropylene glycol, among others, and mixtures thereof. Especially suitable glycols are ethylene glycol and diethylene glycol.

The properties of the low free glycol-containing polyester polyol of the present invention fall within rather broad ranges. The viscosities (Brookfield) of the polyol measured in cps. at 25 °C fall within a rather broad range from about 500 to about 500,000, preferably about 500 to about 100,000, and most preferably about 2,000 to about 50,000; the hydroxyl number values fall within a range of from about 115 to about 750, preferably about 150 to about 600, and most preferably from about 160 to about 320; and the acid number falls within a range of from about .2 to about 40 and preferably about .2 to about 10.

Highly preferred low free glycol-containing polyester polyols of the invention are derived from the reaction of phthalic anhydride or phthalic acid with ethylene and/or diethylene glycol, and are characterized by a viscosity in cps at 25°C of about 8,000 to 35,000, a free glycol content of from about 0 through 5 percent by weight of the polyester polyol, and a hydroxyl number within a range of from about 190 to 240.

The polymer foams of the present invention can be prepared by using standard techniques known to those skilled in the art. In an advantageous embodiment of the invention, the low free glycol-containing polyester polyols are employed in admixture with at least one other polyol in the production of the foams, especially the polyurethane foams. The polyisocyanurate and polyurethane foams can be prepared by mixing together the quasi-prepolymer and the low free glycol-containing polyester polyol, catalyst and blowing agent at temperatures ranging from about 0 °C to 150 °C.

The polyisocyanurate or polyurethane foam of the invention particularly comprises the reaction product of an isocyanate group terminated quasi-prepolymer and a polyol component which comprises

(a) a polyester polyol having a low free glycol content or
(b) a polyol mixture of the low free glycol-containing polyester polyol with at least one other polyol, in the
presence of a blowing agent, a catalyst, and auxiliaries and additives as required. The amount of the low
free glycol-containing polyester polyol in polyol mixture (b) is advantageously sufficient to reduce the aged
k-factor of the resultant foam to below the aged k-factor of the corresponding foam prepared from the same
foam-forming composition and having the same index except that the polyol component solely consists of
the other polyol or polyols and excludes the low free glycol-containing polyester polyol. Preferably, the
polyol component employed in manufacturing the polyisocyanurate or polyurethane foam of the invention
comprises

(a) a polyester polyol having a free glycol content of less than about 7 percent by weight of the polyester polyol or (b) a mixture of the free glycol-containing polyester polyol with at least one other polyol, the total free glycol content of polyol mixture (b) being less than about 7 percent by weight of the content of polyester polyol in said mixture.

The polyurethane foams can be prepared by reacting the polyol of the invention and quasi-prepolymer on an essentially 1:1 to 1:1.25 equivalent basis. In the embodiment wherein the low free glycol-containing polyester polyols of the invention are combined with another polyol(s) to produce polyurethane foams, the low free glycol-containing polyester polyols of the invention can comprise about 5 to 100, preferably about 20 to 80, and more preferably about 40 to 60, weight percent of the total polyol content in the foam preparations.

The polyisocyanurate foams of the invention are prepared by reacting the quasi-prepolymer with a minor amount of polyol, such as sufficient polyol to provide about 0.10 to 0.70 hydroxyl equivalents of polyol per equivalent of said polyisocyanate prepolymer, wherein the low free glycol-containing polyester polyol of the invention comprises about 5 to 100, and preferably about 50 to 100, weight percent of the total polyol content in the foam preparations.

In order to ensure complete reaction in the production of the polyisocyanurate foams, the quasiprepolymer and the polyol are generally mixed in an equivalent ratio of 1.5:1 to 12:1 and preferably 2:1 to 10:1. In ranges outside these proportions the reaction tends to yield a product having less desirable physical charact ristics.

The polyols which can b employed in combination with the low free glycol-containing polyester polyols of the invention in the preparation of the polyurethane and polyisocyanurate foam compositions include

monomeric polyols and polyeth r polyols in general. The polyether polyols ar found particularly useful in preparing rigid polyur thane foams. The polyfunctional active hydrogen initiators used in producing the polyether polyols preferably have a functionality of 2-8, and more preferably have a functionality of 3 or greater (e.g., 4-8).

Any suitable blowing agent can be employed in the foam compositions of the present invention. Water, air, nitrogen, carbon dioxide, readily volatile organic substances and/or compounds which decompose to liberate gases (e.g., azo compounds may be used). Typically, these blowing agents are liquids having a boiling point between minus 50°C and plus 100°C and preferably between 0°C and 50°C. The preferred liquids are hydrocarbons or halohydrocarbons. Examples of suitable blowing agents include, among others, chlorinated and fluorinated hydrocarbons such as trichlorofluoromethane, CCl₂FCClF₂, CCl₂FCFC₂, CClFCClF₂, trifluorochloropropane, difluorodichloromethane, 1-fluoro-1,1-dichloroethane, 1,1-trifluoro-2,2-dichloroethane, 1,1-difluoro-1-chloroethane, methylene chloride, diethylether, isopropyl ether, n-pentane, cyclopentane, 2-methylbutane, methyl formate, carbon dioxide and mixtures thereof. Trichlorofluoromethane is a preferred blowing agent.

The foams also can be produced using a froth-foaming method, such as the one disclosed in U. S. Patent No. 4,572,865. In this method, the frothing agent can be any material which is inert to the reactive ingredients and easily vaporized at atmospheric pressure. The frothing agent advantageously has an atmospheric boiling point of -50° to 10° C, and includes carbon dioxide, dichlorodifluoromethane, monochlorodifluoromethane, trifluoromethane, monochlorotrifluoromethane, monochloropentafluoroethane, vinyl-fluoride, vinylidenefluoride, 1,1-difluoroethane, 1,1,1-trichlorodifluoroethane, and the like. Particularly preferred is dichlorodifluoromethane. A higher boiling blowing agent is desirably used in conjunction with the frothing agent. The blowing agent is a gaseous material at the reaction temperature and advantageously has an atmospheric boiling point ranging from about 10 to 80° C. Suitable blowing agents include trichloromonofluoromethane, 1,1,2-trichloro-1,2,2-trifluoroethane, acetone, pentane, and the like, preferably trichloromonofluoromethane.

The foaming agents, e.g., trichlorofluoromethane blowing agent or combined trichlorofluoromethane blowing agent and dichlorodifluoromethane frothing agent, are employed in an amount sufficient to give the resultant foam the desired bulk density which is generally between 0.5 and 10, preferably between 1 and 5, and most preferably between 1.5 and 2.5, pounds per cubic foot (between 8.0 and 160.0, preferably between 16.0 and 80.0, and most preferably between 24.0 and 40, kilograms per cubic meter). The foaming agents generally comprise from 1 to 30, and preferably comprise from 5 to 20 weight percent of the composition. When a foaming agent has a boiling point at or below ambient, it is maintained under pressure until mixed with the other components. Alternatively, it can be maintained at subambient temperatures until mixed with the other components. Mixtures of foaming agents can be employed.

Any suitable surfactant can be employed in the foams of this invention. Successful results have been obtained with silicone/ethylene oxide/propylene oxide copolymers as surfactants. Examples of surfactants useful in the present invention include, among others, polydimethylsiloxane-polyoxyalkylene block copolymers available from the Union Carbide Corporation under the trade names "L-5420" and "L-5340" and from the Dow Coming Corporation under the trade name "DC-193". Other suitable surfactants are those described in U.S. Pat. Nos. 4,365,024 and 4,529,745 and supplied by Sloss Industries Corporation under the trademarks Foamstab 100 and 200. Generally, the surfactant comprises from about 0.05 to 10, and preferably from 0.1 to 8, weight percent of the foam-forming composition.

Any of the catalysts conventionally employed in the art to catalyze the reaction of an isocyanate with an isocyanate-reactive compound can be employed in the foam preparations of the invention. Such catalysts include organic and inorganic acid salts of, and organometallic derivatives of, bismuth, lead, tin, iron, antimony, uranium, cadmium, cobalt, thorium, aluminum, mercury, zinc, nickel, cerium, molybdenum, vanadium, copper, manganese, and zirconium, as well as phosphines and tertiary organic amines. Examples of such catalysts are dibutyltin dilaurate, dibutyltin diacetate, stannous octoate, lead octoate, cobalt maphthenate, triethylamine, triethylenediamine, N,N,N,N, tetramethylethylenediamine, 1,1,3,3-tetramethylguanidine, N, N, N,N, tetramethyl-1,3-butanediamine, N,N-dimethylethanolamine, N, N-diethylethanolamine, and the like. The catalysts generally comprise from about 1 to 20, and preferably from 0.3 to 10, weight percent of the total foam-forming composition.

In the preparation of the polyisocyanurate rigid foams, any catalysts known to catalyze the trimerization of isocyanates to form isocyanurates, and to catalyze the reaction of isocyanate groups with hydroxyl groups to form polyurethanes, can be employed. One preferred type of catalyst is a mixture of a tertiary amino phenol, such as 2.4.6-tris(dimethylamino methyl)phenol (sold by Rohm and Haas Co. under th designation "DMP-30"), and an alkall metal carboxylate, such as potassium-2- thyl hexoat, the synthesis and use of which are described in U.S. Patent No. 4,169,921. The equivalent ratio of tertiary amino phenol

to alkali metal carboxylate in the cocatalyst composition is desirably about .4:1 to 2.5:1. Another pr ferr d catalyst system is a mixture comprising (i) a salt of a low molecular weight carboxylic acid selected from the group consisting of an alkali metal salt and an alkaline earth metal salt and mixtures thereof (e.g., potassium acetate), (ii) a salt of a higher molecular weight carboxylic acid selected from the group consisting of an alkali metal salt and an alkaline earth metal salt and mixtures thereof, the higher molecular weight carboxylic acid having from about 5 to 30 carbon atoms (e.g. potassium octoate), and (iii) a tertiary amine (e.g., 2,4,6-tris [dimethylaminomethyl] phenol). This mixture is described in U. S. Patent No. 4.710.521.

Other additives may also be included in the foam formulations. Included are processing aids, viscosity reducers, such as 1-methyl-2-pyrrolidinone, nonreactive and reactive flame retardants, such as tris(2-chloroethyl)-phosphate, dispersing agents, plasticizers, mold release agents, antioxidants, compatibility agents and fillers and pigments. The use of such additives is well known to those skilled in the art.

The present invention also provides a process for producing a laminate which comprises (a) contacting at least one facing sheet with a foam-forming mixture comprising the quasi-prepolymer, polyol composition, blowing agent, catalyst, and auxiliaries and additives as required (e.g., a surfactant), and (b) foaming the foam-forming mixture. The process is advantageously conducted in a continuous manner by depositing the foam-forming mixture on a facing sheet being conveyed along a production line, and preferably placing another facing sheet on the deposited mixture. The foam-forming mixture is conveniently thermally cured at a temperature from about 20 °C to 150 °C in a suitable apparatus, such as an oven or heated mold.

Any facing sheet previously employed to produce building panels can be employed in the present invention. Examples of suitable facing sheets include, among others, those of kraft paper, aluminum, and asphalt impregnated felts, as well as laminates of two or more of the above.

The foam materials of the invention can also be used, with or without a facer(s), for pipe insulation.

The foam materials of the invention can contain various reinforcement materials, such as a quantity of glass fibers, as described in U.S. Patent Nos. 4,118,533 and 4,284,683.

The invention is further illustrated by the following examples in which all parts and percentages are by weight unless otherwise indicated.

EXAMPLE 1

This example illustrates the preparation of polyisocyanurate foams (18% trimer) separately using a polyisocyanate and a polyisocyanate prepolymer in combination with polyester polyols having different free glycol contents.

In each foam preparation, the quantities of ingredients listed below in Table I were added to a vessel. First, the A-component (a mixture of the isocyanate, blowing agent and surfactant) was blended in the vessel and cooled to 15.6° C. The B-component (polyester polyol) at 25° C was then added to the vessel, and all ingredients were mixed at 3600 rpm for 10 seconds. The C-component (catalyst) at 25° C was next mixed into the contents of the vessel over a 2-second interval. All ingredients were thereafter mixed at 3600 rpm for an additional 10 seconds and then poured into a box, yielding a rigid polyisocyanurate foam.

Characteristics of the foams produced (Foams A-D) are shown in Table I. The data shown in the table demonstrate that foams prepared from the quasi-prepolymer (Foams C and D) have lower thermal conductivity measurements than foams prepared from an unmodified isocyanate (Foams A and B). The data also show that the thermal conductivity of foam made from the unmodified isocyanate can be lowered by using a low free glycol-containing polyester polyol (Foam B) in place of a polyester polyol containing a higher free glycol content (Foam A). The optimum thermal conductivity lowering is attained by using the quasi-prepolymer in conjunction with the low free glycol-containing polyester polyol in foam production (Foam D).

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TABLE I

	PREPARATION OF POLYISOCYANURATE FOAMS							
5		FOAM						
•	INGREDIENTS(g)	A	В	С	D			
	A-Component							
10	Isocyanate ¹ Isocyanate ² R-11B Surfactant ³	464.0 110.0 4.0	432.0 110.0 4.0	520.4 114.0 4.0	497.4 114.0 4.0			
	B-Component							
15	Polyol PS-3152C ⁴ Polyol PS-2002 ⁵	136.0	168.0	79.6 	102.6			
	C-Component							
20	Catalyst ⁶ Reactivity, sec	23.4 17/26/33/35	23.0 15/25/29/33	23.2 16/-/46/50	23.1 13/37/45/70			
	(Cream/Gel/Firm/Tack-free) Density ⁷ , ib/ft ³ % Closed Cells	1.78 87.6	1.88 89.1	1.79 89.5	1.83 89.7			
25	k-Factor Aging of 1" Thick Core Samples ⁸ (Btu-in/hr-ft ² - F)							
	initial 11 days 29 days	.117 .133 .142	.112 .124 .129	.117 .130 .132	.108 .118 .121			
30	60 days 90 days 120 days	.148 .152 .154	.136 .140 .143	.140 .144 .146	.128 .130			
35	182 days 277 days 364 days Ak-Factor, 364 days	.158 .162 .184 .047	.148 .152 .156 .044	.152 156 .158 .041	.135 .140 .142 .034			

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Footnotes:

- 1. Isocyanate having an equivalent weight of 138, an acidity of 0.03% HCl, and a viscosity of 2000 cps at 25°C.
 - 2. Polyester polyol derived quasi-prepolymer having a viscosity of 13,650 cps at 25° C and a free isocyanate content of 24.8 % (BASF Corporation).
 - 3. Surfactant supplied by the Union Carbide Corporation under the trade name L-5340.
- 4. Polyol PS-3152C = reaction product of phthalic anhydride and diethylene glycol (DEG) having a hydroxyl number of 320, a viscosity at 25°C of 2,500 cps and 15% free DEG (Stepan Company).
 - 5. Polyol PS-2002 = reaction product of phthalic anhydride and DEG having a hydroxyl number of 200, a viscosity at 25 °C of 9,120 cps and 4% free DEG (Stepan Company).
 - 6. Catalyst = mixture employed in the form of a solution in DEG in a weight ratio of 1.18 potassium acetate:1.62 potassium octoate:0.69 DMP-30: 6.51 DEG.
 - 7. $lb/ft^3 \times 16.0 = kg/m^3$.
 - 8. Btu-in/hr-ft² $^{\circ}$ F x .144 = watts/m- $^{\circ}$ C.

EXAMPLE 2

This example illustrates the synthesis of polyisocyanurate foams (10% trimer) using a quasi-prepolym r and a low free glycol-containing polyester polyol.

The foams were prepared according to the procedure of Example 1 from the ingredients and quantities thereof shown in the following Table II. Characteristics of the foams produced are shown in the table. The Table II results show the significant improvement in insulating properties brought about through use of a quasi-prepolymer and a low free glycol-containing polyester polyol, and especially through the combined use of the isocyanate prepolymer and polyester polyol.

TABLE II

	PREPARATION OF POLYISOCYANURATE FOAMS						
_	FISCHARA	FOAM					
15	INGREDIENTS(g)	A	В	С	D		
	A-Component						
20	isocyanate¹ isocyanate² R-11B Surfactant³	380.0 104.0 4.0	326.0 104.0 4.0	418.0 108.0 4.0	375.2 108.0 4.0		
	B-Component	 		181.2			
25	Polyol PS-3152C ⁴ Polyol PS-2002 ⁵	220.0	174.0	161.2	224.8		
	C-Component						
30	Catalyst ⁶ Reactivity, sec	19.4 20/33/40/40	19.6 13/22/28/29	19.5 19/36/47/50	19.4 10/24/33/33		
	(Cream/Gel/Firm/Tack-free) Density ⁷ , lb/ft ³ % Closed Cells	1.68 89.1	1.56 90.8	1.59 86.3	1.65 86.1		
35	k-Factor Aging of 1" Thick Core Samples ⁸ (Btu-in/hr-ft ² - F)				.112		
	Initial 11 days 29 days	.130 .138 .146	.114 .124 .128	.127 .135 .142 .148	.112 .116 .120		
40	60 days 95 days 120 days	.152 .153 .159	.138 .142 .145	.152 .152 .152	.124 .128 .134		
45	180 days 270 days 384 days	.160 .165 .167 .037	.150 .152 .038	.160 .163 .036	.136 .138 .026		

^{1.} Of Example 1.

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7.
$$lb/ft^3 \times 16.0 = kg/m^3$$
.

^{2.}

^{3. &}quot;

^{4. *}

^{5.} Of Example 1.

^{6. &}quot;

^{8.} Btu-in/hr-ft² - *F x .144 = watts/m-* C.

EXAMPLE 3

This example illustrates the preparation of a quasi-prepolymer and its use in combination with polyester polyols having different free glycol contents in the production of polyisocyanurate foam (10% trimer).

A. Preparation of Quasi-Prepolymer

A dry reaction vessel equipped with an agitator, thermometer, addition funnel and inlet for nitrogen gas was charged with 90.1 parts of a polyisocyanate. The isocyanate was agitated under a nitrogen blanket throughout the preparation. The polyisocyanate in the vessel was heated to 90 ± 2° C. 9.9 parts of a polyester polyol? were added at a constant rate over a period of 90 to 120 minutes maintaining the temperature at 90 ± 2° C. After the polyol addition was completed, the reaction was continued at 90 ± 2° C for an additional 90 minutes. The quasi-prepolymer produced had a viscosity of 13,750 cps at 25° C and a free isocyanate content of 26.39%.

- 1. Isocyanate having an equivalent weight of 133 and a viscosity of 180-200 cps at 25 °C.
- 2. Polyol = reaction product of phthalic anhydride and ethylene glycol having a hydroxyl number of 300 and a viscosity at 25 °C of 31,600 cps.

B. Preparation of Polyisocyanurate Foams

The foam syntheses were conducted according to the procedure of Example 1 utilizing the ingredients and quantities thereof presented in Table III below. Characteristics of the foams produced (Foams A-C) are shown in the table. The data show that the combined use of the quasi-prepolymer and low free glycol-containing polyester polyol results in a lower thermal conductivity than use of the quasi-prepolymer with a polyester polyol having a higher glycol content.

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TABLE III

	PREPARATION OF POLYISOCYANURATE FOAMS						
_	11007100100		FOAM				
5	INGREDIENTS	Α	В	С			
	(pts. by wt.)						
	A-Component						
10	Isocyanate ¹ Isocyanate ² R-11B Surfactant ³	253 65.0 2.67	269 56.8 2.67	242 62.4 2.67			
15	B-Component						
	Polyol PS-3152C ⁴ Polyol PS-2002 ⁵	147	131	158			
	C-Component						
20	Catalyst ⁵ Density ⁷ , lb/ft ³ This is Care Samples (Btusin/hr-ft ² F)	13.3	13.3 1.68	13.3 1.72			
	k-Factor Aging of 1" Thick Core Samples ⁸ (Btu-in/hr-ft ² - F)	.128	.118	.106			
25	10 days	.137	.128	.114			
	30 days 60 days 90 days	.150 .154	.136 .140	.118 .122			
30	 Of Example 1. Quasi-prepolymer of Example 3A. Of Example 1. Of Example 1. Of Example 1. 						
35	6. Of Example 1. 7. lb/ft³ x 16.0 = kg/m³. 8. Btu-in/hr-ft² - ° F x .144 = watts/m-° C.						

EXAMPLE 4

This example illustrates the synthesis of further polyisocyanurate foams (18% trimer) utilizing a quasiprepolymer in combination with polyester polyols having different free glycol contents.

The foams were prepared according to the procedure of Example 1 utilizing the ingredients and quantities thereof presented in Table IV below. The data in the table again reveal the improved insulating properties resulting from use of a quasi-prepolymer in combination with a low free glycol-containing polyester polyol in foam production.

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TABLE IV

	PREPARATION OF POLYISOCYANURATE FOAMS						
5		FOAM					
•	INGREDIENTS	Α	В	С			
	(pts. by wt.)						
10	A-Component						
10	Isocyanate¹ Isocyanate² R-11B Surfactant³	309 73.0 2.67	331 59.2 2.67	319 65.4 2.67			
15	B-Component						
	Polyol PS-3152C ⁴ Polyol PS-2002 ⁵	91 	69 	81			
	C-Component						
20	Catalyst ⁶ Density ⁷ , lb/ft ³ k-Factor Aging of 1° Thick Core Samples ⁸ (Btu-in/hr-ft ² - °F)	13.3 1.72	13.3 1.77	13.3			
25	Initial 10 days 30 days 60 days 90 days	.122 .133 .141 .147 .154	.125 .130 .132 .140 .150	.112 .117 .122 .130 .132			
30	 Of Example 1. Quasi-prepolymer of Example 3A. Of Example 1. Of Example 1. Of Example 1. Of Example 1. 						
35	6. Of Example 1. 7. lb/ft ³ x 16.0 = kg/m ³ . 8. Btu-in/hr-ft ² - ° F x .144 = watts/m-° C.						

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Claims

- 1. A polyisocyanurate foam comprising the reaction product in the presence of a blowing agent and
 - (a) an isocyanate group-terminated quasi-prepolymer prepared by reacting an organic polyisocyanate with an organic compound having at least two active hydrogen atoms, and
 - (b) a minor-amount of polyol component, characterized in that the polyol component comprises
 - (1) a polyester polyol having a free glycol content of less that about 7 percent by weight of the polyester polyol or
 - (ii) a mixture of the low free glycol-containing polyester polyol with at least one other polyol, the total free glycol content of mixture (b) being less that about 7 percent by weight of the polyester polyol.
 - 2. The polylsocyanurate foam of Claim 1, characterized in that quasi-prepolymer (a) is the reaction
- product of 55 (a) th organic polyisocyanate, the organic polyisocyanate preferably being toluene diisocyanate, diphenylmethane diisocyanate, polymeric MDI, or mixtures thereof and
 - (b) an aromatic polyester polyol, the aromatic polyester polyol preferably being the reaction product of
 - (i) either (a) est r-containing by-products from the manufacture of dimethyl terephthalat , (b) scrap

polyalkylen ter phthalates, (c) phthalic anhydride, (d) residues from the manufactur of phthalic acid or phthalic anhydrid, () ter phthalic acid, (f) residues from the manufactur of terephthalic acid, (g) isophthalic acid, (h) trimellitic anhydride, (i) phthalic acid, or (j) combinations thereof, and

(ii) at least one low molecular weight aliphatic polyhydric alcohol,

- and polyol component (b) is an aromatic polyester polyol, the aromatic polyester polyol preferably being the reaction product of
 - (i) either (a) ester-containing by-products from the manufacture of dimethyl terephthalate, (b) scrap polyalkylene terephthalates, (c) phthalic anhydride, (d) residues from the manufacture of phthalic acid of phthalic anhydride, (e) terephthalic acid, (f) residues from the manufacture of terephthalic acid, (g) isophthalic acid, (h) trimellitic anhydride, (i) phthalic acid, or (j) combinations thereof, and

(ii) at least one glycol or a mixture of polyols containing at least one glycol.

- 3. The polyisocyanurate foam of Claims 1 or 2, characterized in that quasi-prepolymer (a) is the reaction product of
- (a) either toluene diisocyanate, diphenylmethane diisocyanate, polymeric MDI, or mixtures thereof, the organic polyisocyanate preferably being polymeric MDI, and
- (b) an aromatic polyester polyol, the aromatic polyester polyol preferably being the reaction product of
 - (i) phthalic anhydride and
 - (ii) at least one low molecular weight aliphatic polyhydric alcohol.
- 4. The polyisocyanurate foam of Claims 1,2 or 3, characterized in that quasi-prepolymer (a) is reacted with an aromatic polyester polyol which is the reaction product of a polycarboxylic acid component, the polycarboxylic acid component preferably being phthalic anhydride, phthalic acid, or mixtures thereof, and an aliphatic diol of the generic formula:

HO-R-OH

wherein R is:

- (a) an alkylene radical containing from 2 through 6 carbon atoms,
- (b) a radical of the formula:

-(R10)m-R1-

wherein R1 is an alkylene radical containing from 2 through 6 carbon atoms, and m is an integer of from 1 30 through 4, or

(c) mixtures thereof,

the aliphatic diol preferably being ethylene glycol, diethylene glycol, or mixtures thereof.

- 5. The polyisocyanurate foam of Claims 1, 2, 3, or 4, characterized in that quasi-prepolymer (a) is the reaction product of the organic polyisocyanate and an aromatic polyester polyol utilized in an amount of from 2 to 30 percent by weight relative to the total quasi-prepolymer weight, the aromatic polyester polyol preferably being characterized by a viscosity in cps at 25°C of about 8,000 to 35,000, a free glycol content of from about 0 through 5 percent by weight of the polyester polyol, and a hydroxyl number within the range of from about 190 to 240.
- 6. The polyisocyanurate foam of any one of the preceding claims, characterized in that quasiprepolymer (a) is reacted with an aromatic polyester polyol having a free glycol content of less than about 5 percent by weight of the aromatic polyester polyol.
 - 7. The polyisocyanurate foam of any one of the preceding claims, characterized in that the blowing agent is water, air, nitrogen, carbon dioxide, readily volatile organic substances, compounds which decompose to liberate gases, or mixtures thereof, and preferably is a chlorinated or fluorinated hydrocarbon, mixtures of said hydrocarbons, or mixtures of said hydrocarbons with water, and the foam optionally includes a viscosity reducer, flame retardant, dispersing agent, plasticizer, mold release agent, antioxidant, compatibility agent, filler or pigment, or mixtures thereof.
 - 8. A laminate comprising at least one facing sheet adhered to the polyisocyanurate foam of any one of the preceding claims 1 to 7.
 - 9. A polyurethane foam comprising the reaction product in the presence of a blowing agent and a catalyst of
 - (a) an isocyanate group-terminated quasi-prepolymer prepared by reacting an organic polyisocyanate with an organic compound having at least two active hydrogen atoms, and
 - (b) a polyol component, characterized in that the polyol component comprises a polyol mixture containing a polyester polyol having a free glycol content of less than about 7 percent by w ight of th polyester polyol, the amount of the low free glycol-containing polyester polyol in the polyol mixture being sufficient to reduc the ag d k-factor of the resultant foam to below the aged k-factor of the corresponding foam pr pared from the same foam-forming composition and having the same index except that the polyol

component does not include the low fr e glycol-containing polyester polyol.

10. A laminate comprising at least one facing sheet adhered to the polyur than foam of claim 9.